Molecular dynamics simulations of momentum and thermal diffusion properties of near-critical argon along isobars

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A B S T R A C T

Three basic diffusion properties of argon – shear viscosity, bulk viscosity and thermal conductivity – were studied in the neighborhood of the critical point using molecular dynamics (MD) and the Lennard-Jones potential energy function. MD simulations were performed along the 1.0P and 1.2P, isobars. Green-Kubo relations and a Lennard-Jones pair potential were used. Four different sets of Lennard-Jones parameters were used. A comparison of computed shear viscosity and thermal conductivity values with data available from the National Institute of Standards and Technology (NIST) displayed a good agreement. Results for bulk viscosity indicated that values of this property cannot be neglected in this thermodynamic region, a result that violates the traditional and much-assumed Stokes hypothesis in classical fluid mechanics. Furthermore, it was shown that in the neighborhood of the critical region the bulk viscosity can have larger values than the shear viscosity.

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1. Introduction

Fluid dynamics and heat transfer models are strongly related to accurate values for transport coefficients. The Navier-Stokes equation [1] and the energy equation [2] incorporate for a given substance three transport coefficients that connect the physical dynamics of momentum and heat to three properties, namely, bulk viscosity, shear viscosity and thermal conductivity.

There are in the literature several data sets that collect experimental values of shear and thermal conductivity for simple substances. The electronic tables from the National Institute of Standards and Technology (NIST) [3], for instance, allow the solution of a great number of typical engineering problems. In these cases, the bulk viscosity is generally neglected based on the validity of Stokes hypothesis [4] even when the fluid is neither incompressible nor a monatomic dilute gas [5]. Hence, bulk viscosity values are rarely found in tables, technical works and engineering projects, even though they are needed for the complete specification of the Navier–Stokes stress tensor.

However, in the neighborhood of the critical point, availability of transport coefficient data is scarce. Moreover, theory and experiment display anomalies in the behavior of some thermodynamic and transport properties, usually appears as a divergence towards infinite or zero depending on the property [6–10]. In particular, the high compressibility of near-critical fluids reduces the accuracy of experimental measurements. For this class of fluids, bulk viscosity data are even more limited since the acoustic absorption of sound waves – the most used experimental technique [11,12] – is subject to high levels of error. To overcome this drawback, Brillouin scattering was developed as a promising technique to measure bulk viscosity, though it is still not widely known and used [13].

In order to overcome the aforementioned experimental limitations, an interesting possibility is performing molecular simulations that can suitably model the near-critical state and compute transport coefficients. One of these methods is equilibrium molecular dynamics (MD) [14–16]. In this approach, fundamental thermophysical properties such as energy, temperature, pressure can be determined from specific averages of the dynamic evolution of an ensemble of molecules. Other properties are obtained using relations from thermodynamics and statistical mechanics including time correlations, e.g., transport properties [17]. This method is useful in the investigations of phase diagrams [18] and chemical reactions [19]. In the framework of MD, particle tra-
jectories in phase space defined by positions and velocities are computed from the forces resulting from an interaction potential energy between pairs of atoms and molecules. The most employed model for non-bonded particles is the Lennard-Jones (L-J) potential [20]. It has a simple mathematical expression and leads to accurate results notably in the study of noble gases. The L-J has already been used to compute transport properties of supercritical fluids far from their critical points, transport properties of near-critical fluids using other approaches and other properties [21–26].

In this work, we investigated transport properties of near-critical argon using equilibrium MD. The present simulations are based on Green-Kubo theory and use four different versions of argon Lennard-Jones potentials available in the literature. The importance of investigating argon in this work is twofold. First, it is the most abundant noble gas in the atmosphere. Second, it is an important substance employed for calibration purposes in laboratories because it is chemically inert and has a low cost. The present results were compared with available experimental data, which allows one to estimate their range of applicability. Due to the scarcity of bulk viscosity experimental values, special emphasis was given to this transport property in such a way that they can be used in simulations of compressible fluid flow [27] as well as heat transfer problems [28].

2. Fundamentals

2.1. Molecular dynamics and the Lennard-Jones potential

The basis of the MD technique is the computational simulation of Newton’s equation of motion for a set of interacting moving particles. For a system consisting of argon atoms in a regular box, only interactions between pairs of non-bonded atoms were considered. In other words, many-body interactions are not considered explicitly.

A pair potential suitably models the interactions between atom pairs. The simplest realistic model for this purpose is the L-J 12-6 potential. It is a two parameters model given by Eq. (1)

\[
U_{ij}(r_{ij}) = \begin{cases} 
4\epsilon \left[ \frac{\sigma}{r_{ij}} \right]^{12} - \frac{\sigma}{r_{ij}} \right]^{6}, & r_{ij} \leq \sigma_c \\
0, & r_{ij} > \sigma_c 
\end{cases}
\]

(1)

where \( \epsilon \) is the depth of the potential well, \( \sigma \) is the zero-energy distance, \( r_{ij} \) is the distance between particles \( i \) and \( j \), and \( \sigma_c \) is a cut-off distance that plays an important role in reducing the computational cost. To use this potential it is necessary to correct the computed value of the potential in order to include the small energies resulting from long-range interactions farther than \( \sigma_c \) [20]. The 12-exponent term models repulsion effects while the 6-exponent
term describes attraction effects. Each i–j pair of particles interacts via a force \( \mathbf{f}_{ij} \) that can be calculated from
\[
\mathbf{f}_{ij} = -\nabla U_{LJ}(r_{ij})
\] (2)

The motion of each particle is determined from its resultant force. An important feature of MD is that an accurate potential energy expression generally leads to accurate thermophysical properties. When using the Lennard-Jones 12-6 expression, all possible model choices are restricted to selecting \( \sigma \) and \( \varepsilon \) values in addition to a special care concerning the \( r_c \) values in all cases. In order to consider uncertainties of the LJ potential parametrization of argon, four different pairs of \( \sigma \) and \( \varepsilon \) and their respective critical constants were used and compared in this work. They are shown in Table 1.

In MD simulations it is possible to adopt the concept of statistical ensemble from statistical mechanics. One can simulate an
isolated system (e.g., microcanonical or NVE ensemble), a closed system (canonical, isochoric-isothermal or NVT ensemble), an isobaric-isothermal system (NPT ensemble), among other possibilities, without using partition functions that may be difficult to compute. Before physical arguments are considered, the choice of an ensemble is related to the use of a specific numerical integration scheme of the equations of motion. Each algorithm can include artificial thermostats or barostats to control the basic thermodynamic variables—N, V, T and P [14]. The mathematical expressions used in MD and derived from statistical mechanics theory for computing the thermophysical properties of interest must be obtained from suitable ensembles.

It is also important to avoid edge effects. For this purpose, MD simulations of bulk systems generally employ periodic boundary conditions. These artificial boundaries imposed on the system restrict the simulation box size. This approach is called minimal-image convention and it states that the characteristic dimension of the simulation box—the length for a cubic box—must be greater than twice the cutoff radius \( r_c \).

2.2. Green-Kubo formulas for transport properties

Equilibrium MD employs Green-Kubo formulas [31] for the computation of shear and bulk viscosity as well as of thermal conductivity. The three coefficients could be obtained experimentally by perturbing the system towards non-equilibrium states and observing its response that is associated with irreversible processes. Therefore, in principle, to compute them one should use non-equilibrium molecular dynamics (NEMD) instead of equilibrium MD. NEMD is based on the direct application of a heat or momentum flux on the simulation box [32,33]. On the other hand, the fluctuation dissipation theorem allows the computation of transport coefficients from the microscopic fluctuations of the system. This theorem states that small fluctuations of the system are related to internal relaxation processes. Therefore, transport coefficients then can be calculated as integrals of time-correlation functions of the microscopic stress tensor and internal energy fluxes in an equilibrium state.

Shear viscosity \( \eta_s \) is related to the resistance of the fluid to shear forces. In the microcanonical and canonical ensembles this

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**Table 1**

Parameter pairs for the Lennard-Jones 12-6 potential argon atoms along with their respective critical constants.

<table>
<thead>
<tr>
<th>Model Parameters</th>
<th>Critical Constants</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \sigma (\text{Å}) )</td>
<td>( \kappa/\kappa_0 (\text{K}) )</td>
<td>( T_c (\text{K}) )</td>
</tr>
<tr>
<td>3.405</td>
<td>119.8</td>
<td>150.86</td>
</tr>
<tr>
<td>3.43</td>
<td>119.3</td>
<td>150.86</td>
</tr>
<tr>
<td>3.345</td>
<td>125.7</td>
<td>150.86</td>
</tr>
<tr>
<td>3.3952</td>
<td>116.79</td>
<td>153.00</td>
</tr>
</tbody>
</table>

\( T_c \): critical temperature; \( P_c \): critical pressure; \( \rho_c \): critical density; \( \kappa_0 \): Boltzmann constant.
property is obtained by integrating the ensemble average of the time correlation function of the stress tensor $P$ off-diagonal elements, which represent the momentum fluxes,

$$n_s = \frac{1}{3k_BT} \sum_{m<n} \int_0^\infty \langle \sigma_{mn}(0)\sigma_{nm}(t) \rangle dt$$

(3)
Fig. 7. Bulk viscosity of $10^4$ argon atoms for 1.0P, isobar in a range of reduced temperatures from 1.0 to 2.0 using Lennard-Jones model with different parameters pairs – (a) [25], (b) [26], (c) [29], (d) [30] – and (e) the average values of $\eta_b$ compared to the average values of $\eta_b$ for the same isobar.

The ensemble average of the time correlation function of the stress tensor $\mathbf{J}^p$ diagonal elements minus the product $P \nu$. In other words,

$$\eta_b = \frac{1}{3k_BVT} \sum_j \int_0^\infty (\langle J^p_{mn}(0) - P(0)\nu \rangle J^p_{mn}(t) - P(t)\nu) dt$$

(4)

Thermal conductivity, $\kappa$, physically expresses the ability of the substance in transferring thermal energy when subject to temperature gradients. It is computed by integrating the ensemble average of time correlation function of the microscopic heat flux $\mathbf{J}^Q$ vector, namely,

$$\kappa = \frac{1}{3k_BVT} \int_0^\infty \langle J^Q(0) \cdot J^Q(t) \rangle dt$$

(5)

In Eqs. (3) and (4), the stress tensor $\mathbf{J}^p$ can be computed from

$$\mathbf{J}^p = \sum_{i=1}^N m_i \mathbf{v}_i \mathbf{v}_i - \sum_{i=1}^N \sum_{j>i}^N \mathbf{r}_{ij} \cdot (\nabla U_{ij})$$

(6)

where $m_i$ is the mass of the particle $i$, $U_{ij}$ the interaction potential between particles $i$ and $j$, and the prime means matrix transpose. Diagonal elements have the index $m$ equal to $n$. In Eq. (5), the energy flux vector is calculated from

$$\mathbf{J}^Q = \frac{1}{V} \left[ \sum_{i=1}^N e_i \mathbf{v}_i + \frac{1}{2} \sum_{i<j}^N (-\nabla U_{ij} \cdot (\mathbf{v}_i + \mathbf{v}_j)) \mathbf{r}_{ij} \right]$$

(7)

where $e_i$ is the energy, potential and kinetic, of particle $i$ whereas $\mathbf{v}_i$ is the velocity of particle $i$.

2.3. Near-critical effects

The thermophysical and rheological properties of a substance in the neighborhood of the critical point are characterized by an anomalous behavior. These critical anomalies are observed as divergences either to zero or infinity. Theory and experiment show that bulk and shear viscosities and thermal conductivity diverge to infinity in the vicinity of the critical point $[34,23,35–39,21,40,41]$. One important feature of these divergences is their universality, i.e., they are described by universal parameters and expressions that do not depend on the nature of the fluid. These models are power laws having a single parameter that indicates the separation of each thermodynamic state from the critical point. For the two viscosities and thermal conductivity, this parameter is the temperature difference $\Delta T = T - T_c$. However, it is also important to identify the correct thermodynamic path of the power law model, since
different paths lead to different power law coefficients that could be related through an equation of state [6]. For instance, along the critical isochore, one has

\[ \kappa \approx \kappa^\circ \Delta T^\nu + \eta - \gamma \]  
\[ \eta_s \approx \eta_s^\circ \Delta T^{-\eta} \]  

where \( \nu = 0.630 \pm 0.001 \), \( \eta = 0.031 \pm 0.004 \) and \( \gamma = 1.239 \pm 0.002 \) are universal exponents obtained from renormalization group theory [42], with \( \kappa^\circ \) and \( \eta_s^\circ \) depending on the substance. In particular, studies seeking to understand the behavior of bulk viscosity near critical conditions are rare. Five of the most representative ones based either on Monte Carlo simulations, or on MD and even purely theoretical [21,43–46] did not describe the divergence of

**Fig. 8.** Bulk viscosity of 10⁴ argon atoms for 1.2Pc isobar in a range of reduced temperatures from 1.0 to 2.0 using Lennard-Jones model with different parameters pairs – (a) [25], (b) [26], (c) [29], (d) [30] – and (e) the average values of \( \eta_s \) compared to the average values of \( \eta_t \) for the same isobar.

**Fig. 9.** Correlations of diagonal stress tensor elements of 10⁴ argon atoms of 1.0Pc and 1.2Pc, isobars for three temperatures 1.00Tc, 1.10Tc, and 2.00Tc, using Lennard-Jones potential parameters from [26].
this property in terms of the classical critical exponents. Furthermore, scaling laws similar to the ones given by Eqs. (8) and (9) were not found along isobars. Therefore, the present work is especially important since it can help overcoming these difficulties as shown next.

3. Computational methods

The MD simulations were carried out using the open-source code LAMMPS [47]. Shear viscosity and thermal conductivity were computed in the NVT ensemble using a Nosé-Hoover style non-Hamiltonian equation of motion [48] in order to improve the control of the simulation temperature. In order to prevent any influence over pressure fluctuations, bulk viscosity values were obtained in the NVE ensemble using the Verlet algorithm.

Following recent papers focusing on near critical MD simulations [5,49], we used $10^4$ argon atoms in all simulations. This number of particles leads to a better control of the thermodynamic state in the simulation box and improved statistical averages. The numerical integration of the time correlation functions of the transport coefficients (Eqs. (3)–(5)) used the trapezoidal rule over all time steps. Each simulation was equilibrated for 2 ns ($2 \times 10^6$ time steps) before averages were computed for a time span of 10 ns ($10^7$ time steps). A timestep of 1 fs was used. Correlations of heat and momentum fluxes were computed for a sampling length of 200 ps ($2 \times 10^5$ time steps). A large cut-off radius of 8r was chosen to improve the accuracy in the calculation of the interaction energies and better describe long-range correlations typical of the critical region. The correction in the potential for $r_g > r_c$ was computed integrating a unitary radial distribution function for these distances [20]. By employing a large value of the $r_c$ distance, simulations were more realistic for a dense fluid and described more accurately long-range corrections. Periodic boundary conditions were used. A temperature range from 1.0$T_c$ up to 2.0$T_c$ along two isobars – 1.0$P_c$ and 1.2$P_c$ – was considered. The four pairs of L-J parameters shown in Table 1 were used in the computations. More details about the simulation procedure can be found elsewhere [18].

4. Results and discussion

Figs. 1 and 2 show shear viscosity results of argon obtained from the MD simulations along the 1.0$P_c$ and 1.2$P_c$ isobars using four different sets of L-J parameters taken from the literature (Table 1).

There are no significant differences between the results computed using all the L-J parameter pairs according to Table 1. Therefore, computation of the shear viscosity is barely sensitive to small variations in the parameters of the interaction potential. Furthermore, MD simulations were able to reproduce the behavior of the shear viscosity critical divergence quite well. Figs. 1e and 2e show that simulations are in good agreement with NIST data. The results obtained using Michels et al.’s [25] parameters for the L-J potential exhibit a better description of the critical divergence in the shear viscosity.

To ensure the convergence of the results in Figs. 1 and 2, we present in Fig. 3 a set of normalized $F_{nm}$ correlation functions of three representative temperatures for both isobars. The results agree with data already reported in literature [50].

Figs. 4–6 show the computed thermal conductivity values and the heat flux correlation functions of argon, as done for the shear viscosity. The results are also in good agreement with the NIST data. All parameters pairs lead to similar results with small discrepancies as compared to the average values. The parameter pair from Michels et al. [25] displays a smoother behavior in comparison to NIST data and better represented the critical divergence. The error bars are smaller as compared to the shear viscosity ones. In Fig. 6, overall the heat flux correlation function converges more slowly than the respective shear stress correlation in agreement with previously reported information [51]. It is important to point out that this plot shows correlations up to 100 ps and the simulations considered the correlations up to 200 ps.

Finally, Figs. 7–9 collects MD bulk viscosity results for argon and the set of correlation functions used to compute them. Since these data are not available in NIST tables, they can be especially useful. Once again, all four pairs of L-J parameters lead to similar results, but in this case the L-J parameters from White [29] showed a clearly smoother behavior and a better description of the critical divergence. Large deviations were found in the average value of bulk viscosity results. These deviations are considerably larger than deviations reported for shear viscosity and thermal conductivity. Moreover, Fig. 9 shows that the diagonal stress tensor correlations exhibit a better convergence if compared to thermal conductivity correlations. The correlation analysis of the bulk viscosity suggests that this property is easier to compute as compared to thermal conductivity even near the critical point.

Another important feature in Figs. 7 and 8 is the non-zero value of the bulk viscosity far from the critical temperature. At $T_s = 2.0$ (≈300 K) this property reaches values around 30 $\mu$Pa s with an average deviation of ± 5 $\mu$Pa s considering the results obtained from all set of L-J parameters. This result shows that even though argon is a monoatomic gas, the Stokes hypothesis as well as the kinetic gas theory assumptions [52] do not hold and thus cannot be used. Furthermore, this property has the same order of magnitude of the shear viscosity values and, hence, cannot be neglected in simulations of the momentum conservation equations.

Figs. 7e and 8e compare both viscosities obtained using averages over the MD results and the four sets of L-J parameters at each $T_s$ value. These results indicate that bulk viscosity values are actually larger than their shear counterparts for the thermodynamic values considered here. This is an important finding because classical fluid mechanics analysis neglects the bulk viscosity [53]. From these figures we propose an empirical relation between both viscosities at each isobar: one should use $\eta_b \approx 1.3\eta_s$ at the 1.0$P_c$ isobar and $\eta_b \approx 1.6\eta_s$ at the 1.2$P_c$ isobar. Therefore, the traditional assumption of neglecting this difference cannot be a priori assumed in the neighborhood of the critical point.

5. Conclusions

In this work it was shown that molecular dynamics simulations using the Lennard-Jones potential to describe the potential energy between argon atoms allow one to obtain accurate values of both shear viscosity and thermal conductivity along isobars in the neighborhood of the critical point. The computed properties were in good agreement with experimental data collected in the NIST database. This good agreement leads us to compute also the bulk viscosity in the same thermodynamic region even though NIST data is not available in this case.

In the neighborhood of the critical point, the results for bulk viscosity obtained with molecular dynamics show that; (a) shear and bulk viscosities diverge along the same isobar with similar behavior; (b) bulk viscosity does not go to zero away from the critical point in spite of being close to it, thus contradicting Stokes hypothesis and kinetic theory even for a monoatomic dilute gas; (c) bulk viscosity is not only of the same order of magnitude as shear viscosity near the critical point, but it is actually larger than its counterpart in this region.

The present approach of using molecular dynamics to compute diffusion properties of a fluid in the neighborhood of the critical point is general and we have shown that can be used to obtain reliable results in this experimentally difficult to measure
thermodynamic region. The crucial point in the method is the choice of the interaction potential, which can be an issue for more complicated and practical systems (e.g., N₂). Investigations along these lines are under way in our group.

The complete set of results for the transport properties is available as an excel file in the Supplementary material of this work.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.supflu.2016.04.004.

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